600930027 FINAL

DATA EVALUATION REPORT

OIL OF CITRONELLA

Study Type: Mutagenicity: <u>Salmonella typhimurium</u>/Mammalian Microsome Mutagenicity Assay

Prepared for:

Health Effects Division
Office of Pesticide Programs
Environmental Protection Agency
1921 Jefferson Davis Highway
Arlington, VA 22202

Prepared by:

Clement International Corporation 9300 Lee Highway Fairfax, VA 22031-1207

Principal Reviewer J. Habe	Date	5/4/92
Lynne T. Haber, Ph.O.		
Independent Reviewer Warm G. War For	Date	5/4/92
Nancy E. McCarroll, H.S.		, , ,
QA/QC Manager \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	Date	<u>5/4/92</u>
Sharon Segal, Ph.D.		, ,

Contract Number: 68D10075
Work Assignment Number: 1-76

Clement Number: 91-247

Project Officer: James Scott

GUIDELINE SERIES 84: MUTAGENICITY

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EPA Reviewer: J. Thomas McClintock, Ph.D.
Reregistration Section, Science Analysis and

Coordination Branch (H7509C)

EPA Section Head: <u>Albin Kocialski, Ph.D.</u> Reregistration Section, Science Analysis and

Coordination Branch (H7509C)

Signature.

Date:

Signature:

Date: 5/8/97

DATA EVALUATION REPORT

STUDY TYPE: Mutagenicity: Salmonella typhimurium/mammalian microsome

mutagenicity assay

EPA IDENTIFICATION Numbers:

Tox Chem. Number: 21901

MRID Number: 421513-10

TEST MATERIAL: TREO SPF 15 lotion

SYNONYMS: Oil of citronella

SPONSOR: Primavera Laboratories, Inc., New York, NY

STUDY NUMBER: 063629-2

TESTING FACILITY: United States Testing Company, Inc., Hoboken, NJ

TITLE OF REPORT: Ames Salmonella Microsome Mutagenesis Assay on TREO SPF 15

Lotion

AUTHOR: Wang, X.M.

REPORT ISSUED: December 2, 1991

<u>CONCLUSIONS--EXECUTIVE SUMMARY</u>: No conclusions can be reached from the <u>Salmonella typhimurium</u>/mammalian microsome plate incorporation assay conducted with TREO SPF 15 lotion. The study was seriously compromised for the following reasons:

- 1--Unacceptably high spontaneous reversion counts for strains TA98 and TA100.
- 2--Marked reductions in the sensitivity of strains TA98 and TA100 to direct-acting mutagens.
- 3--The use of an excessive concentration of the S9-activated positive control (10 μ g/plate 2-anthramine).

(See Section C, Reported Results, for a detailed discussion.)

Based on the above considerations, it was concluded that the study was unacceptable, and, therefore, did not satisfy Guideline requirements (§84-2) for genetic effects Category I, Gene Mutations.

STUDY CLASSIFICATION: Unacceptable. The study should be repeated using established procedures for the Salmonella typhimurium/mammalian microsome plate incorporation assay.1 The purity of the test compound should also be included with the report of the repeat study.

A. MATERIALS:

1	Test	Material:	TREO	SPF	15	lotion
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Description: Off-white emulsion

Lot number: Not reported Purity: Not reported

Receipt date: Not reported Stability: Not reported Contaminants: None listed

Solvent used: Dimethyl sulfoxide (DMSO)

Other provided information: The storage conditions and frequency of

dose solution preparation were not reported.

2. Control Materials:

Negative: Not done

Solvent/final concentration: DMSO/≤0.1 mL

	Positive:	
	Nonactivation:	10 μg/plate TA1535, TA100
	N-Methyl-N-nitro-	рудаес ппрэзэ, ппры
	N-nitrosoguanidine (MNNG)	50_ μg/plate TA1538, TA98
	2-Nitrofluorene (2-NF)	100 μg/plate TA1537
	9-Aminoacridine (9-AA)	100 µg/place iniss/
	Activation:	
	2-Anthramine (2-AA)	10 μg/plate all strains
3.	Activation: S9 derived from adult male	xratxliver
	x Aroclor 1254 x induced noninduced	mouse lung
		hamster other
	none	other
	other	- Other

¹Maron, D.M. and Ames, B.N. (1983). Revised Methods for the <u>Salmonella</u> Mutagenicity Test. <u>Mutat</u>. <u>Res</u>. 113:173-215.

Neither the strain of rats nor the source of the S9 were reported. S9 mix composition: Concentration/mL Component: 100 µmoles Sodium phosphate buffer (pH 7.4) 5 umoles Glucose 6-phosphate 4 µmoles NADP 33 µmoles KC1 8 umoles MgCl₂ 100 µL S9 4. Test Organism Used: S. typhimurium strains _____ TA97 <u>x</u> TA98 <u>x</u> TA100 ____ TA102 ____ TA104 x TA1535 x TA1537 x TA1538 list any others: Test organisms were properly maintained: Overnight cultures were prepared from master plates held at 4°C for no more than 1 month. is some ambiguity about the source of cultures for the master plates. The study author stated that "master plates are prepared from frozen permanents kept at 0°C," but also stated that cultures are stored "in liquid nitrogen at approximately -190°C." Checked for appropriate genetic markers (rfa mutation, R factor): Yes. 5. Test Compound Concentrations Used: (a) Preliminary cytotoxicity assay: Up to 10,000 μg/plate; the concentrations tested were not specified. (b) Mutation assay: Five doses (50, 150, 500, 1500, and 5,000 μ g/plate) were evaluated in triplicate in the presence and absence of S9 activation; all tester strains were used. B. TEST PERFORMANCE: 1. Type of Salmonella Assay: x Standard plate test _____ Pre-incubation (_____) minutes _____ "Prival" modification ____ Spot test ____ Other (describe) 2. Methods:

the preliminary cytotoxicity assay was not specified. For the mutation assay, 0.1 mL of a 16 ± 4 -hour culture of the appropriate tester strain, and up to $100~\mu\text{L}$ of the appropriate test

(a) Preliminary cytotoxicity/mutation assays: The procedure used for

material dose, solvent, or positive controls were added to tubes containing 2-mL volumes of molten top agar. For the S9-activated tests, 0.5 mL of the S9-cofactor mix was added. Tester strains, and test and control solutions were added as described. The contents of the tubes were mixed, poured over Vogel-Bonner minimal medium E plates, and incubated at 37°C for 48±4 hours. Triplicate plates per strain, per dose, per condition were used for the test compound, solvent, and positive controls.

- 3. <u>Evaluation Criteria</u>: The test material was considered positive for a particular strain and condition if it caused at least a doubling in the number of revertants of strains TA98 or TA100, or a tripling in the number of revertants of strains TA1535, TA1537, or TA1538.
- 4. Statistical Analysis: The data were not analyzed statistically.
- 5. Protocol: See Appendix B.

C. REPORTED RESULTS

- 1. Preliminary Cytotoxicity Assay: Neither the data nor the details of the preliminary cytotoxicity assay were reported. The study author stated that the test material was evaluated in strains TA100 and TA1537 at concentrations up to $10,000~\mu\text{g/plate}$. The study author further stated that cytotoxicity was observed between 1000 and $10,000~\mu\text{g/plate}$. Accordingly, the dose range selected for the mutation assay was 50-5,000 $\mu\text{g/plate}$ +/-S9.
- 2. Mutation Assay: Representative results from the mutation assay with TREO SPF 15 lotion are presented in Table 1. As shown, solvent control colony counts for strain TA98 +S9 and strain TA100 +/-S9 were well beyond the expected range reported by Maron and Ames² (TA98:30-50 revertants/plate; TA100: 120-200 revertants/plate) and either approached the acceptability limit for TA98 (15-75) or exceeded the range for TA100 (60-220) established by other investigators.³ The results with the nonactivated positive controls (10 $\mu g/plate$ MNNG and 50 $\mu g/plate$ 2-NF) further indicated that the higher-than-expected background counts for TA98 and TA100 coincided with a reduction in sensitivity.

Table 1 also shows that there was a marked difference in the number of revertant colonies of TA1538 and TA98 induced by 50 μ g/plate 2-NF. Exposure of these strains to comparable doses of 2-NF generally results in the production of approximately equivalent numbers of revertant colonies. While the response (i.e., fold increase) varies as a function of spontaneous mutant counts, the actual number of mutant colonies induced by 2-NF is consistently close. Similarly, marked differences

²Maron, D.M. and Ames, B.N. (1983).

³deSerres, F.J. and Shelby, M.D. 1979. "Recommendations on Data Production and Analysis Using the Salmonella/Microsome Mutagenicity Assay." <u>Environmental Mutagenesis</u> 1:87-92.

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between the number of mutant colonies induced in strains TA1535 and TA100 by 10 $\mu g/plate$ MNNG were observed in this study, even though a similar phenomenon exists for these strains.

Excessive levels of 2-AA (10 μ g/plate) were used for the S9-activated positive control. Such high levels can be expected to be cytotoxic, resulting in reduced revertant levels. Finally, the reporting of colony counts >3000/plate is an unacceptable practice. The generally accepted rule, whether counting manually or using an automatic colony counter, is that counts \geq 3000 colonies/plate can not be accurately determined.

Based on the above considerations, it was concluded that the study is invalid.

- D. <u>REVIEWERS' DISCUSSION/CONCLUSIONS</u>: The numerous technical deficiencies identified in the review of the study precludes an evaluation of the relevance of the findings with the test material. The study should be repeated using established procedures for the <u>S. typhimurium/mammalian</u> microsome mutagen-icity assay.⁴ Additionally, information about the purity of the test material should be submitted with the repeat study.
- E. <u>QUALITY ASSURANCE MEASURES</u>: Was the test performed under GLP? <u>Yes</u>. (A quality assurance statement was signed but not dated.)
- F. <u>CBI APPENDICES</u>: Appendix A, Introduction and Materials and Methods, CBI pp. 7-9; Appendix B, Protocol, CBI pp. 20-23.

<u>CORE CLASSIFICATION</u>: Unacceptable. The study does not satisfy the data Guideline requirement (§84-2) for genetic effects Category I, Gene Mutations.

⁴Maron, D.M. and Ames, B.N. (1983).

TABLE 1: Representative Results of the <u>Salmonella typhimurium</u>/Mammalian Microsome Mutation Assay with TREO SPF 15 Lotion

	Dose/Plate	S9 Activation	Revertants per Plate of Bacterial Tester Straina				
Substance			TA1535	TA1537	TA1538	TA98	TA100
Solvent Control			<u> </u>				
Dimethyl sulfoxide	≤100 µL	-	19±6	16±3	29±4	43±2	275±16
•	≤100 µL	+	37±4b	13±2	31±7	74±8	370±6
Positive Controls					•		
N-methyl-N-nitro- N-nitrosoguanidine	10 µg	·	4019±113			· - -	1027±123
9-Aminoacridine	100 µg	-	See See	641±405			÷ -
2-Nitrofluorene	50 μg	-			3044±154	1544±169	Same and
2-Anthramine	10 µg	+	469±201	1151±13	3549±44	3753±192	3270±103
Cest Material							
TREO SPF 15 lotion	500 μg ^c	-	25±5	15±6	30±7	47±5	275±12
	1500 µg	-	18±9	7±2	19±7	36±4	309±46
	5000 μg	-	12±3	0±0	10±5	33±3	286±28
	150 µg°	+	28±1	24±3	24±6	109±4	370±23
	500 μg	+	26±11	20±6	28±3	96±13	328±16
	1500 μg	+	28±1	20±6	30±3	90±13	343±32
	5000 μg	+	22±5	20±2	32±10	75±3	306±18

 $^{^{\}mathrm{a}}\mathrm{Means}$ and standard deviations of the counts from triplicate plates.

bCalculated by our reviewers; erroneously reported by the study author as 27±21.

 $^{^{\}circ}$ Results for lower doses (50 and 150 μ g/plate -S9, and 50 μ g/plate +S9) did not suggest a mutagenic effect.

APPENDIX A

INTRODUCTION AND MATERIALS AND METHODS CBI $pp.\ 7\mbox{-}9$

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